Reviewed By: Henry Spencer, Ph.D. Section VII. Toxicology Secondary Reviewer: Albin Kocialski, Ph.D. Section VII, Toxicology Branch (TS-769C) Section VII, Toxicology Branch (TS-769C)

006939

DATA EVALUATION REPORT

Study: Teratology and Postnatal Studies on Picloram in the Rat

Laboratory: Dow Chemical Company; Midland, Michigan

In Fd. Cosmet. Toxicology 10:6, 797-803 (1972)

Study No.: None provided

MRID No.: None provided (000 30 284)

Material Tested: Picloram, Lot No. 2RS17 (purity not stated)

Animals: Sprague-Dawley Rats

Methods:

Four groups of 35 female rats (10 for postnatal portion and 25 for teratology portion) were gavaged with picloram suspended in corn oil on days 6 to 15 of gestation. Dosages were 0, 500, 750, and 1000 mg/kg/day. The females were observed for signs of toxicity on a daily basis. Body weights were obtained on either day 20 or 21 postpartum for teratology. Normal teratology parameters, i.e., live and dead fetuses, resorptions, fetal weights, and skeletal and visceral evaluations were determined. The postnatal evaluations determined the following duration of qestation: litter weights at days 0, 5, and 21, and viability.

Results:

Maternal toxicity was noted at 750 and 1000 mg/kg as hyperactivity and mild diarrhea. Cnly evidence of unossified sternebrae was considered as the fetotoxicity end point (no NOEL). Bilateral accessory ribs were increased in numbers of fetuses, but not by litter, at 1000 mg/kg. There was a lack of evidence for postnatal effects from picloram.

Conclusions:

A fetotoxic NOEL was not determined at 500 mg/kg (LDT)

Toxicity Category: N/A

Core Rating: Supplementary for the lack of a NOEL

Repairability: None

DATA EVALUATION RECORD

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BRANCH TOX DISC	Teratology		
FORMULATION	Technical Material		
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APPROVED BY: TITLE: ORG: LOC/TEL: SIGNATURE:		DATE: 10/11/8	
CONCLUSIONS: Teratology - rat			
1. This is a scientif	icallume lid stude		
2- Picloram is not ter	ratogenic to rats at maternal doses 1000 mg/kg/day doses are toxic to	of 500 to 1000 mg/l	kg/day, S•
uerayed ossiticatio	y toxic to the fetus (judging from on of 5th sternebrae which is usual of 500 to 1000 mg/kg/day. No NOEL	v accordated with a	nstamal .
4. Neonatal developmer at doses of 500 to	nt/survival is not altered by materi 1000 mg/kg/day given on days 6-15 (nally administered of gestation.	picloram
This study is classified for law 100 FL of	as core minimum fortenat lagrans	end suppleme	Tack a
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MATERIALS AND METHODS:

Picloram

Picloram (Lot No. 2RS17) was suspended in corn oil at a concentration of 200 mg/ml for administration to pregnant rats on days 6 to 15 of gestation. Four groups of 35 Sprague-Dawley rats (25 for teratology and 10 for the postnatal portion of the study) received 0, 500, 750 or 1000 mg picloram/kg/day by gavage. Rats were observed daily for signs of toxicity. Pre-breeding and gestation day 20 body weights were obtained on teratology rats and prebreeding and postpartum day 21 body weights were obtained for postnatal females.

The following observations were made:

Fetal Examinations (Teratology Litters)

in utero position

| live fetuses
| complete and partial resorptions
| corpora lutea
| fetal weights
| skeletal exam 2/3 each litter
| visceral exam 1/2 each litter
| (0, 750 and 1000 mg/kg groups
| only)

Postnatal Litters

duration of gestation
labor, delivery date (if possible)
litter size, pup wt at birth and
days 5, 21
sex at birth, day 21
age - eyes open
% viability birth, days 5, 21
skeletal exam weanlings (2/sex/group)

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Maternal body weight gains, numbers of implantations, corpora lutea, resorptions, litter size and pup weights were evaluated at the 95% level by a one-way analysis of variance with Scheffe's Multiple Range Test. Fetal abmormalities among litters and fetal populations were analyzed at the 95% level using the 2 x 2 contingency table and chi-square test.

REPORTED RESULTS:

Dosage at 500 mg/kg/day produced no overt signs of toxicity, while rats given 750 or 1000 mg/kg/day developed hyperaesthesia and mild diarrhea after 1-4 days of treatment, and 14 maternal deaths occurred between days 3 and 17 of gestation in these dosage groups. In surviving dams, doses up to and including 1000 mg/kg/day affected neither maternal weight gains, litter size and resorption rate nor other reproductive parameters examined. Evidence of retarded fetal growth, as reflected by an increase in unossified fifth sternebrae was observed in all of the treated groups but not in an exactly dose-related manner. The occurrence of bilateral accessory ribs was increased significantly in fetuses of dams given 1000 mg/kg/day for 10 days during gestation, but in the usual evaluative mode (% litters) there was no significant effect.

DISCUSSION:

This study was conducted in a scientifically appropriate manner.
 Maternally toxic and non-effect doses were used.

- 2. Complete review of this study is possible with the use of the "raw data" package in addition to the written publication.
- 3. Picloram appears to have no effect on the fetus that is unrelated to maternal toxicity -- and then the fetus appears to be fairly well protected from the insult upon the maternal animal.

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